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Wavelets in Numerical Analysis

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WAMA

Cargèse, July 2004

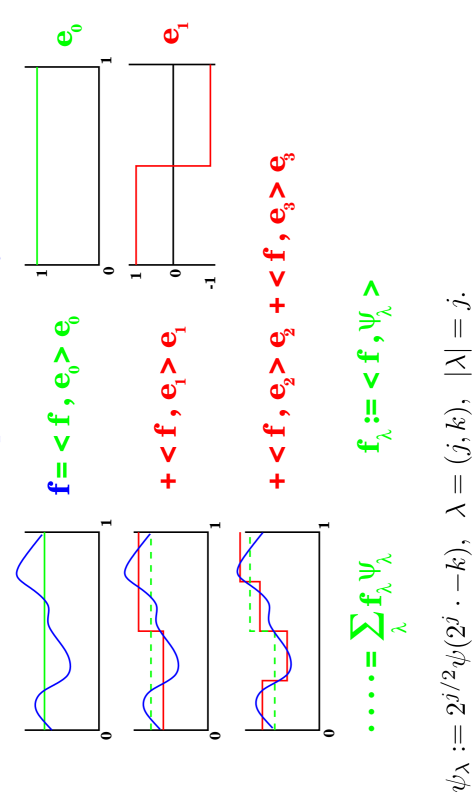
Agenda

1. Motivation: adaptive wavelet discretizations and PDE's

2. Adaptive space refinement for operator equations

3. Adaptive multiresolution processing for evolution equations

Basic example: the Haar system

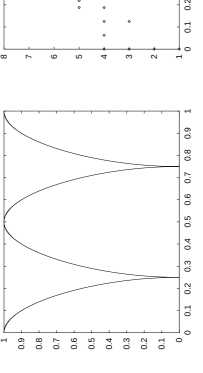


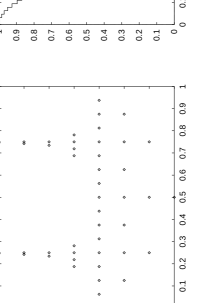
approximation processes, using smoother functions such as splines More general wavelets are constructed from similar multiscale or finite elements.

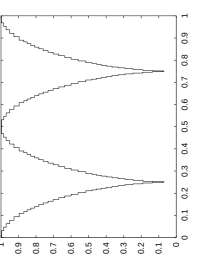
Approximating functions by wavelet bases

- Linear approximation at resolution level j by taking the truncated sum $f \mapsto P_j f := \sum_{|\lambda| \le j} f_{\lambda} \psi_{\lambda}$.
- Nonlinear (adaptive) approximation obtained by thresholding

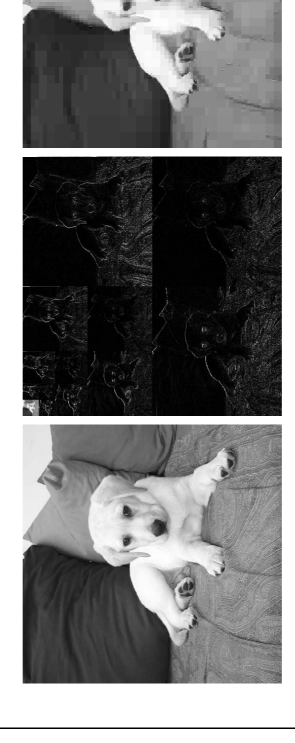
$$f \mapsto \mathcal{I}_{\Lambda} f := \sum_{\lambda \in \Lambda} f_{\lambda} \psi_{\lambda}, \quad \Lambda = \Lambda(\eta) = \{\lambda \text{ s.t. } |f_{\lambda}| \ge \eta\}.$$







Applications to image compression



decomposition 10³ largest coefficients

digital picture

Linear approximation results

- V_h : finite element space discretizing a domain $\Omega \subset \mathbb{R}^d$.

-
$$N := \dim(V_h) \sim \operatorname{vol}(\Omega) h^{-d}$$

-
$$W^{s,p} := \{ f \in L^p(\Omega) \text{ s.t. } D^{\alpha} f \in L^p(\Omega), |\alpha| \le s \}$$

Ciarlet-Raviart, Strang-Fix): provides with the classical estimate Classical finite element approximation theory (Bramble-Hilbert,

$$f \in W^{s+t,p} \Rightarrow \inf_{g \in V_h} ||f - g||_{W^{s,p}} \le Ch^t \sim CN^{-t/d},$$

assuming that V_h has enough polynomial reproduction and is contained in W_p^s .

Nonlinear approximation results

N-terms approximations: $\Sigma_N := \{ \sum_{\lambda \in \Lambda} d_\lambda \psi_\lambda ; \#(\Lambda) \le N \}.$

Rate of decay governed by weaker smoothness conditions (DeVore): with 1/q = 1/p + t/d

$$f \in W^{s+t,q} \Rightarrow \inf_{g \in \Sigma_N} ||f - g||_{W^{s,p}} \le CN^{-t/d},$$

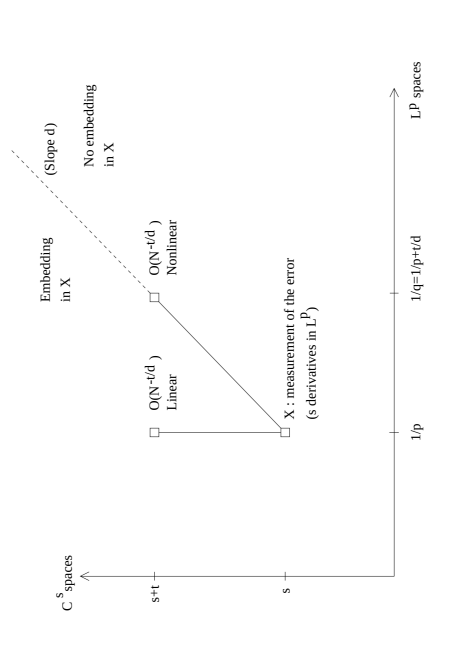
approximation is obtained by thresholding: if $f = \sum_{\lambda} d_{\lambda} \psi_{\lambda}$, and - For most error norm X (e.g. L^p , $W^{s,p}$, $B^s_{p,q}$), a near optimal $f_N := \sum_{N \text{ largest } ||d_{\lambda}\psi_{\lambda}||_X} d_{\lambda}\psi_{\lambda}$, we then have

$$||f - f_N||_X \le C \inf_{g \in \Sigma_N} ||f - g||_X$$

with C independent of f and N.

- Remark: a similar theory for piecewise polynomial approximation on N adaptive triangles is still to be completed.

Pictorial interpretation of approximation results



General program for PDE's

- conservation laws (DeVore and Lucier 1987), elliptic problems on certain PDE's might have substantially higher regularity in the - Theoretical: revisit regularity theory for PDE's. Solutions of scale governing nonlinear approximation than in the scale governing linear approximation. Examples: hyperbolic corner domains (Dahlke and DeVore, 1997).
- appropriate adaptive resolution strategies which perform essentially $||u - \tilde{u}_N||$ has the same rate of decay N^{-s} as $||u - u_N||$ in some - Numerical: develop for the unknown u of the PDE $\mathcal{F}(u) = 0$ as well as thresholding : produce \tilde{u}_N with N terms such that prescribed norm, if possible in $\mathcal{O}(N)$ computation.

Two approaches toward adaptive wavelet methods

- formulation of the problem. (Bertoluzza, Perrier, Liandrat, Canuto, appropriate discretization sets $\{\psi_{\lambda}\}_{{\lambda}\in\Lambda_n}$, based on a variational Dahlke, Hochmuth, Urban, Masson, Dahmen, DeVore, AC). - First approach (mostly applied to steady steate problems $\mathcal{F}(u) = 0$): iterative space refinement techniques to access
- computational time and memory size, while preserving the accuracy of the initial scheme (Harten, Abgrall, Arandiga, Chiavassa, Donat, $\partial_t u = \mathcal{E}(u)$: multiresolution adaptive post-processing, i.e. start from a classical and reliable scheme on a uniform grid and use a discrete multiresolution decomposition in order to compress - Second approach (mostly applied to evolution problems Dahmen, Mueller, Farge, Schneider, Kaber, Postel, AC)

General variational problems

solution of $\mathcal{F}(u) = 0$, i.e. $D\mathcal{F}(u)$ is an isomorphism from \mathcal{H} to \mathcal{H}' . \mathcal{H} Hilbert space, $\mathcal{F}: \mathcal{H} \to \mathcal{H}'$ continuous mapping, u nonsingular

Variational formulation: find $u \in \mathcal{H}$ such that

$$\langle \mathcal{F}(u), v \rangle = 0$$

for all $v \in \mathcal{H}$.

Simple linear examples: $\mathcal{F}(u) = \mathcal{A}u - f$

- Laplace: $\mathcal{A} := -\Delta$ and $\mathcal{H} := H_0^1$
- Stokes: $\mathcal{A}(u,p) := (-\Delta u + \nabla p, -\operatorname{Div} u)$ and $\mathcal{H} := (H_0^1)^3 \times L_0^2$.
- Single layer potential $\mathcal{A}u(x) := \int_{\Gamma} \frac{u(y)}{4\pi|x-y|} dy$ and $\mathcal{H} := H^{-1/2}$.

Standard (FEM) approach to discretisation

- 1. Well posed problem in infinite dimension $\mathcal{F}(u) = 0$.
- 2. Finite dimensional discretization $\mathcal{H} \to V_h$ by a Petrov-Galerkin type method $(\langle \mathcal{F}(u_h), v_h \rangle = 0 \text{ for all } v_h \in W_h).$

Difficulties: not always well-posed (compatibility conditions, e.g.

LBB for Stokes: $\inf_{p_h \in P_h} \sup_{u_h \in U_h} \frac{\int p_h \operatorname{Div} u_h}{\|p_h\|_{L^2} \|u_h\|_{H^1}} \ge \beta_h > 0$).

3. Iterative solver $u_h^0 \to u_h^1 \cdots \to u_h$.

Difficulties: ill-conditionning and dense matrices

4. Adaptivity: derive local error indicators by a-posteriori analysis of residual $\mathcal{F}(u_h)$, and apply local mesh refinement based on these indicators $V_h = V_r^0 \to V_r^1 \to \cdots$, $u_h = u_r^0 \to u_r^1 \to \cdots$ Difficulties: hanging nodes, convergence analysis of such refinement strategies (Dörfler 1996, Morin-Nocetto-Siebert 2000). Wavelet adaptive discretizations: new paradigm

- 1. Well posed problem in infinite dimension $\mathcal{F}(u) = 0$.
- 2. Equivalent discrete problem in infinite dimension by wavelet-Galerkin: find $U = (u_{\lambda})_{{\lambda} \in \nabla}$ such that

$$F(U) := (\langle \mathcal{F}(\sum u_{\lambda} \psi_{\lambda}), \psi_{\mu} \rangle)_{\mu \in \nabla} = 0.$$

renormalization, i.e. $||u||_{\mathcal{H}}^2 \sim \sum |u_{\lambda}|^2$ and $||u||_{\mathcal{H}'}^2 \sim \sum |\langle u, \psi_{\lambda} \rangle|^2$. Well-posed: $F: \ell^2 \to \ell^2$ if $(\psi_{\lambda})_{\lambda \in \nabla}$ is a Riesz basis for \mathcal{H} after

- 3. Converging iteration in infinite dimension $U^0 \to U^1 \to \cdots \to U$.
- tolerances in finite dimension: U^n supported by finite wavelet set 4. Adaptive approximation of this iteration up to prescribed

⇒ allows to establish optimal accuracy and complexity results in the energy $||u||_{\mathcal{H}} \sim ||U||$ norm.

The linear elliptic case

Assume \mathcal{A} is an \mathcal{H} -elliptic operator. Equivalent problem:

$$AU = F$$

where A is ℓ^2 -elliptic. For a suitable κ the iteration,

$$U^{n+1} = U^n + \kappa [F - AU^n],$$

Approximate iteration with prescribed tolerance $\varepsilon > 0$,

converges with fixed error reduction rate $\rho < 1$.

$$U^{n+1} = U^n + \kappa[\text{APPROX}(F, \varepsilon) - \text{APPROX}(AU^n, \varepsilon)],$$

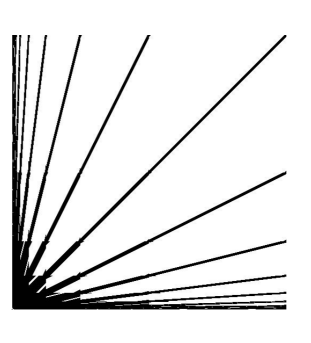
with $\|\text{APPROX}(AU^n, \varepsilon) - AU\| \le \varepsilon$ and $\|\text{APPROX}(F, \varepsilon) - F\| \le \varepsilon$. converges with reduction rate ρ until error is of order ε .

The procedure APPROX (F,ε) amounts in thresolding F in ℓ^2 , or equivalently the data f in the \mathcal{H}' norm.

Matrix-vector approximation

compression: one can build A_N with N coefficients per rows and The procedure APPROX (AU^n, ε) is made possible by matrix column such that $||A - A_N|| \le CN^{-r}$

(W ₃ ,V ₁)	(W_3,W_1)	$(W_3.W_2)$	(W_5,W_5)
(W_2,V_1)	(W_2, W_1)	(W_2,W_2)	(W ₂ ,W ₃)
(W ₁ ,V ₁)	(W ₁ , W ₁)	(V ₁ ,W ₂) (W ₁ ,W ₂)	(V ₁ ,W ₁) (W ₁ ,W ₃)
(V ₁ ,V ₁) (W ₁ ,V ₁)	(V, W,)	(V_1,W_2)	(V, .W,)



Analysis: based on the Schur lemma, using esimates of the type

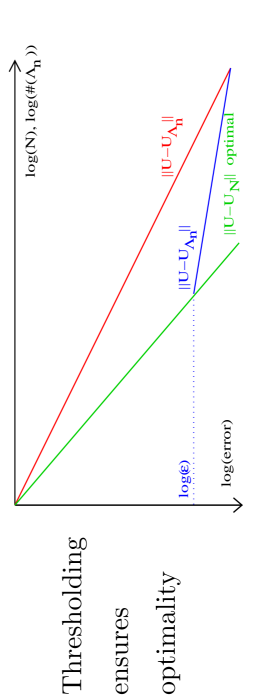
$$|\langle \mathcal{A}\psi_{\lambda}, \psi_{\mu} \rangle| \le C[1 + \operatorname{dist}(\lambda, \mu)]^{-\beta} 2^{-\gamma ||\lambda| - |\mu||},$$

derived from the smoothness and vanishing moments of the ψ_{λ} .

The role of thresholding

 $||V-U|| \le \varepsilon$, then with a > 1 fixed and W the smallest subvector Lemma: if U is such that $||U - U_N|| \le CN^{-s}$ and V is such that of V such that $||V - W|| \le a\varepsilon$, one has

$$||U-W|| \le (1+a)\varepsilon$$
 and $\#(W) \le C\varepsilon^{-1/s}$, i.e. $||U-W|| \le C[\#(W)]^{-s}$



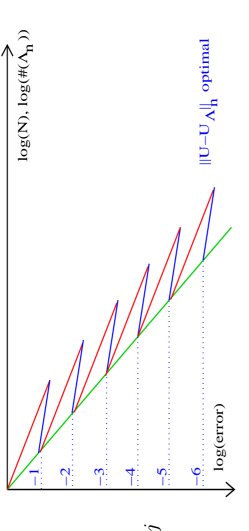
Problem: intermediate memory size and computational time should also be optimal, i.e. $\mathcal{O}(\varepsilon^{-1/s})$.

Geometric tolerances

Idea: decrease

tolerances
$$\varepsilon_0 = 1$$
,

$$\varepsilon_1 = \frac{1}{2}, \cdots, \varepsilon_j = 2^{-j}$$



Fixed number of iteration at each step $j \to j+1$ involving sparse decomposing $V = V_1 + [V_2 - V_1] + [V_4 - V_2] + \cdots$, and taking matrix-vector product: $W = APPROX(AV, \varepsilon)$ obtained by

$$W := A_2 J V_1 + A_2 J_{-1} [V_2 - V_1] + \dots + A_1 [V_2 J_{-1}]$$

with J large enough such that

$$||W - AV|| \le ||A|| ||V - V_{2^J}|| + \sum_{j=1}^J ||A - A_{2^{J-j}}|| ||V_{2^j} - V_{2^{j-1}}|| \le \varepsilon.$$

Results

Theorem (Dahmen, DeVore, AC - Math. Comp. 2000): if V is such that $||V - V_N|| \le CN^{-s}$, and if $||A - A_N|| \le CN^{-r}$ with r>s, then $|\operatorname{Supp}(W)|\leq C\varepsilon^{-1/s}$ and therefore $|W - AU|| \le C|\operatorname{Supp}(W)|^{-s}.$

ingredients (thresholding, adaptive matrix vector multiplication) Theorem (Dahmen, DeVore, AC - FoCM 2002): The general strategy for linear operator equations based on the above achieves the ultimate goal, namely production of U^n and $\Lambda_n = \operatorname{Supp}(U^n)$, such that if $||U - U_N|| \le CN^{-s}$, then

$$||U - U^n|| \le C \# (\Lambda_n)^{-s},$$

with $\mathcal{O}(\#(\Lambda_n))$ computational cost.

Remarks on practical aspects

All wavelet properties are exploited: Sobolev norm equivalences, smoothness (not always available) and vanishing moments.

Coarsening is not needed in all practical cases studied so far, yet Dahmen and DeVore, using the Morin-Nocetto-Siebert algorithm seems necessary in the proof of the optimality theorem! Similar optimality results recently obtained for adaptive FEM by Binev, combined with coarsening. Complexity is dominated by assembling matrix elements, numerical quadratures, addressing the indices in Λ_n (key role of efficient data wavelets based on the same FE spaces: for a given error, wavelets may win for $N_{\rm d.o.f.}$ but lose (by a factor > 4) for computational structures). Practical comparison between adaptive FEM and

Extension to more general problems

Saddle point problems $AU + B^TP = F$ and BU = G, e.g. based on adaptive approximation of the Uzawa iteration (Dahlke, Hochmuth and Urban 1999):

$$AU^{n} = F - B^{T}P^{n-1}$$
 and $P^{n} = P^{n-1} + \kappa(BU^{n} - G)$

adaptive FEM algorithm: Nocetto 2002. Question: do the same No LBB is needed here, adaptivity stabilizes Similar result for $-\varepsilon\Delta u + a.\nabla u = 0$ with convergence rate independent of ε ? concepts apply to convection dominated problems, such as

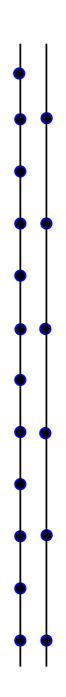
(need specific adaptation of fast evaluation of F(U)), no available Extension to nonlinear problems: DeVore, Dahmen, A.C. 2002 numerical results yet. Problem dependent tuning seems unavoidable in order to optimize this type of algorithms.

A discrete multiresolution framework

- Γ_j , $j = 0, \dots, J$: sequence of discretisations at scales 2^{-j} .
- $U_j = (U_j(\gamma))_{\gamma \in \Gamma_j}$ discretisation of a fonction u on Γ_j , i.e. vector of $\mathcal{V}_j := \mathbb{R}^{\Gamma_j}$.
- Restriction operator P_{j-1}^j from V_j onto V_{j-1} : computes coarser discretization $U_{j-1} = P_{j-1}^j U_j$ from the next finer.

Basic example 1: point values on nested grids $\Gamma_{j-1} \subset \Gamma_j$, i.e.

$$U_{j-1}(\gamma) = U_j(\gamma) \text{ for } \gamma \in \Gamma_{j-1}.$$



Basic example 2: cell averages on nested partitions, i.e.

$$U_{j-1}(\gamma) = \operatorname{vol}(\gamma)^{-1} \sum_{\mu \in \Gamma_j, \mu \subset \gamma} \operatorname{vol}(\mu) U_j(\mu)$$

- Prediction operator P_i^{j-1} from V_{j-1} into V_j : reconstructs an approximation $\hat{U}_j = P_j^{j-1} U_{j-1}$ of U_j .
 - Consistancy assumption: $P_{j-1}^{j}P_{j}^{j-1}=I$

Point value example: $\hat{U}_j(\gamma) = U_{j-1}(\gamma)$ for $\gamma \in \Gamma_{j-1}$, and $\hat{U}_j(\gamma)$ obtained par local interpolation for $\gamma \in \Gamma_j \setminus \Gamma_{j-1}$.



averages in a consistant way, e.g. via polynomial reconstruction. Cell avergage example: $\hat{U}_j(\gamma)$ obtained by "interpolating" the

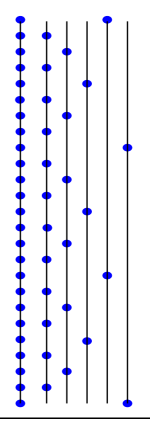


Multiscale decomposition

Prediction error $E_j := U_j - \hat{U}_j \in \mathcal{W}_{j-1} = \operatorname{Ker}(P_{j-1}^j)$. Detail vector D_{j-1} : coordinates of E_j in a basis of W_{j-1} . Point value example: $D_{j-1}(\lambda) = E_j(\lambda), \ \gamma \in \Gamma_j \setminus \Gamma_{j-1}$ interpolation error at intermediate point. Cell average example: on each coarse cell of Γ_{j-1} the prediction error E_j has null average \Rightarrow define D_j by removing for each coarse cell γ one fine cell $\mu \subset \gamma$.

$$U_J \Leftrightarrow (U_{J-1}, D_{J-1}) \Leftrightarrow (U_{J-2}, D_{J-2}, D_{J-1}) \Leftrightarrow \cdots$$

 $\Leftrightarrow (U_0, D_0, \cdots, D_{J-1}) = \mathcal{M}U_J = (d_{\lambda})_{{\lambda} \in \nabla_J}$



Physical grid Γ_J

Multiscale grid (point values) ∇_J

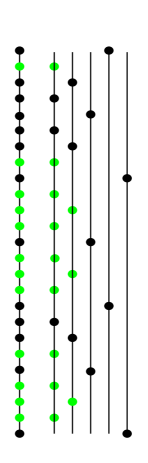
Complexity of \mathcal{M} and \mathcal{M}^{-1} : $\mathcal{O}(\operatorname{Card}(\Gamma_J))$

Compression

Thresholding: given a level dependent threshold $\eta = (\eta_0, \dots, \eta_{J-1})$ set to zero all coefficients $|d_{\lambda}| \leq \eta_{|\lambda|} \Leftrightarrow \text{approximation of } U_J \text{ by}$

$$\mathcal{I}_{\eta}U_J = \mathcal{I}_{\Lambda}U_J = \mathcal{M}^{-1}\mathcal{R}_{\Lambda}\mathcal{M}U_J,$$

 \mathcal{R}_{Λ} : restriction of ∇_J to $\Lambda = \Lambda(\eta) = \{\lambda \in \nabla_J \; \text{ t.q. } |d_{\lambda}| \geq \eta_{|\lambda|} \}$.



Adaptive mesh $\Gamma(\Lambda)$

Adaptive set Λ

cell averages $U_J(\gamma)$ on an adaptive physical mesh $\Gamma(\Lambda)$ associated to Λ , (need to impose that Λ has a tree structure up to enlarging it). Compressed representation $(d_{\lambda})_{{\lambda}\in\Lambda}$ to the data of point values or

Complexity of adaptive decomposition and reconstruction algorithms: $\mathcal{O}(\operatorname{Card}(\Lambda))$

Adaptive multiresolution processing

Reference scheme on Γ_J : approximation of $u(x, n\Delta t)$ by $U_J^n = (U_J^n(\gamma))_{\gamma \in \Gamma_J}$ with $U_J^{n+1} = E_J U_J^n$

$$U_J^{n+1}(\gamma) = U_J^n(\gamma) + F(U_J^n(\mu) \; ; \; \mu \in S(\gamma)).$$

 $S(\gamma)$: local stencil (excludes implicit schemes).

In the case of FV conservative schemes, F has the form of a balance over the edges surrounding the cell γ

$$U_J^{n+1}(\gamma) = U_J^n(\gamma) + \sum_{\mu \text{ s.t. } |\Gamma_{\gamma,\mu}| \neq 0} F_{\gamma,\mu}^n$$

where $F_{\gamma,\mu}^n = -F_{\mu,\gamma}^n$ is a function of the $U_J^n(\nu)$ for ν in a local stencil surrounding γ and μ .

Adaptive algorithm

physical values (point values or cell averages) on the adaptive mesh $V_J^n = (V_J^n(\gamma))_{\gamma \in \Gamma_J}$ is represented by its coefficients $(d_\lambda^n)_{\lambda \in \Lambda_\eta^n}$ or its Goal: compute approximations of $u(x, n\Delta t)$ by (V_J^n, Λ_η^n) , where $(V_J^n(\gamma))_{\gamma\in\Gamma(\Lambda_\eta^n)}$ (we always impose the graded tree structure on

Benchmark: an ideal choice would be Λ_n^n the smallest graded tree adaptive solution V_I^n should still be comparable to $\mathcal{I}_{\eta}U_J^n$, i.e. the corresponding thresolding operator applied to the exact reference containing $\{\lambda, |d_{\lambda}(U_J^n)| \geq \eta_{|\lambda|}\}$ but it is not be accessible. The solution.

Initialization: define Λ_n^0 the smallest graded tree containing $\{\lambda, |d_{\lambda}(U_J^0)| \ge \eta_{|\lambda|}\}$ and set $V_J^0 := \mathcal{I}_{\eta}U_J^0$, Derivation of $(V_J^{n+1}, \Lambda_{n+1})$ from (V_J^n, Λ_n)

Three basic steps:

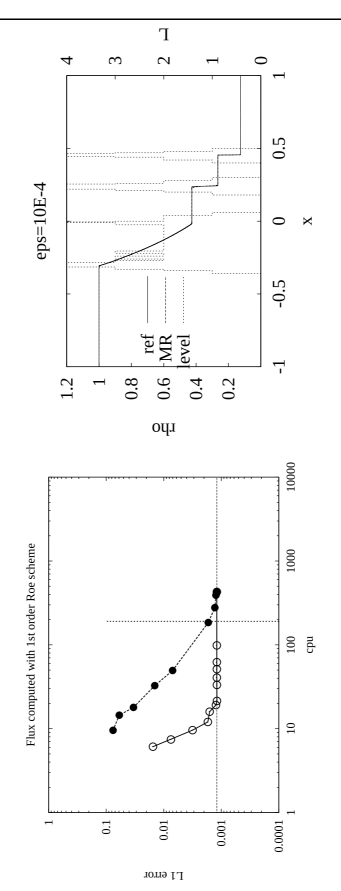
- the solution at time n+1 (ideally such that $|d_{\lambda}(E_J V_J^n)| < \eta_{|\lambda|}$ if - Refinement: predict a superset $\Lambda_{\eta}^{n} \subset \tilde{\Lambda}_{\eta}^{n+1}$ adapted to describe $\lambda \notin \tilde{\Lambda}_{\eta}^{n+1}$) and extend by $d_{\lambda}^{n} = 0$ for $\lambda \in \tilde{\Lambda}_{\eta}^{n+1} \setminus \Lambda_{\eta}^{n}$.
- Evolution: compute the new value $V_J^{n+1}(\gamma)$, for $\gamma \in \Gamma(\tilde{\Lambda}_{\eta}^{n+1})$ (ideally $V_J^{n+1} = \mathcal{I}_{\tilde{\Lambda}_\eta^{n+1}} E_J V_J^n$).
- Coarsening: apply level dependent thresholding operator to the computed vector $(d_{\lambda}^n)_{\lambda \in \tilde{\Lambda}_{\eta}^{n+1}} \Rightarrow \text{new set } \Lambda_{\eta}^{n+1} \subset \tilde{\Lambda}_{\eta}^{n+1} \text{ and } V_J^{n+1}$.

The loss of accuracy with respect to the reference scheme depends on each of the three steps.

- Coarsening: accuracy is controled by the level of the threshold η and the stability properties of the multiscale reconstruction (existence of underlying continuous wavelet systems).
- discrete evolution operator E_J on the size of the coefficients in the - Refinement: accuracy is controled by analyzing the action of the multiscale decomposition.
- Evolution: need an accurate evolution step in the compressed form. Two possible approaches:
- (i) direct application of the numerical scheme on the adaptive grid $\Gamma(\tilde{\Lambda}_{\eta}^{n+1})$: loss of accuracy for low order schemes.
- reconstruction on fine grid: more accurate but more costly. (ii) exact computation of $\mathcal{T}_{\tilde{\Lambda}_n^{n+1}}E_JV_J^n$ by local adaptive

Numerical illustration

In 1D: comparison of AMR and local reconstruction on Sod tube test.



preserves the accuracy with a substantial reduction of CPU time For a low order reference scheme, only local reconstruction and memory space (1/20 at best).

Error Analysis

Remark: adaptive evolution with local reconstruction is given by

$$V_J^{n+1} = \mathcal{I}_{\Lambda_\eta^{n+1}} \mathcal{I}_{\tilde{\Lambda}_\eta^{n+1}} E_J V_J^n.$$

Compare $U_J^{n+1} = E_J U_J^n$ with $V_J^{n+1} = \mathcal{I}_{\Lambda_{n+1}} \mathcal{I}_{\tilde{\Lambda}_{n+1}} E_J V_J^n$.

Cumulative error analysis between both solutions:

$$||U_J^{n+1} - V_J^{n+1}|| \le ||E_J U_J^n - E_J V_J^n|| + d_n,$$

with $d_n = ||V_J^{n+1} - E_J V_J^n|| \le t_n + c_n$ where

$$t_n := \|\mathcal{T}_{\Lambda_{n+1}}\mathcal{T}_{\tilde{\Lambda}_{n+1}}E_JV_J^n - \mathcal{T}_{\tilde{\Lambda}_{n+1}}E_JV_J^n\|, \quad c_n := \|\mathcal{T}_{\tilde{\Lambda}_{n+1}}E_JV_J^n - E_JV_J^n\|,$$

refinement and thresholding strategies should allow to control both denote the thresholding and refinement errors. The analysis of terms with a prescribed precision ε .

Controling the thresholding error

Analysis based on underlying continuous wavelet system (ψ_{λ}) :

$$||U_J - \mathcal{I}_{\Lambda} U_J|| \le \sum_{\lambda \notin \Lambda} ||d_{\lambda} \psi_{\lambda}||.$$

For the L^1 norm, this gives $||U_J - \mathcal{I}_\Lambda U_J|| \le C \sum_{\lambda \notin \Lambda} 2^{-d|\lambda|} |d_\lambda|$, and therefore with $\eta_j = 2^{dj}\eta_0$,

$$||U_J - \mathcal{I}_{\eta} U_J|| \le C \sum_{2^{-d|\lambda|}|d_{\lambda}| < \eta_0} 2^{-d|\lambda|} |d_{\lambda}|$$

- Crudest estimate: $\eta_0 \#(\nabla_J) \sim \eta_0 2^{dJ} \Rightarrow \text{take } \eta_0 = \varepsilon 2^{-dJ}$.
- Better estimate: $\eta_0 \#(\tilde{\Lambda}^{n+1}) \Rightarrow \text{take } \eta_0 = \varepsilon / \#(\tilde{\Lambda}^{n+1})$.
- Even better: take largest η_0 s.t. $\sum_{2-d|\lambda||d\lambda|<\eta_0} 2^{-d|\lambda|}|d\lambda| \le \varepsilon$.

Controling the refinement error

Harten's refinement rule for hyperbolic equations (assuming CFL condition for the reference scheme $\Delta t \leq C2^{-J}$): - If $|d_{\lambda}| > \eta_{|\lambda|}$ include in $\tilde{\Lambda}_{\eta}^{n+1}$ the neighbors of λ at the same level.

- If $|d_{\lambda}| > 2^{r-1}\eta_{|\lambda|}$ also include the childrens of λ at the finer level.

Here r represents the order of accuracy of the prediction operator. Not sufficient to prove that $|d_{\lambda}(E_J V_J^n)| < \eta_{|\lambda|}$ if $\lambda \notin \tilde{\Lambda}_{\eta}^{n+1}$.

This can be proved by a more severe refinement rule: refine of nlevel if $2^{n(s-1)}\eta_{|\lambda|} \leq |d_{\lambda}| < 2^{(n+1)(s-1)}\eta_{|\lambda|}$, with s the Hölder smoothness of the underlying wavelet system. In practice, however, we observe that Harten's rule is sufficient and that the thresolding error dominates the refinement error.

A crude error estimate

scheme in the sense that $||E_JU - E_JV|| \le (1 + c\Delta t)||U - V||$, this Assuming stability in the prescribed norm $\|\cdot\|$ for the reference yields the cumulative (too pessimistic) estimate

$$||U_J^{n+1} - V_J^{n+1}|| \le (1 + c\Delta t)||U_J^n - V_J^n|| + \varepsilon \le \dots \le C(T)n\varepsilon \sim \frac{\varepsilon}{\Delta x}$$

Main defects of this analysis:

- In most practical cases, we observe that thresholding and refinement error does not accumulate linearly.
- the relevant Besov-Sobolev smoothness which governs the nonlinear coefficients which is used to represent the adaptive solution (we are not ensured that $N \ll \#(\Gamma_J)$. Such a bound would require that - No error bound available in terms of the number N. of wavelet approximation error in $\|\cdot\|$ is preserved under the action of evolution and thresholding.